

**IN THE UNITED STATES PATENT AND TRADEMARK OFFICE**

Applicants : Mouritsen et al.  
Serial No. : 08/955,373  
Filed : October 21, 1997  
Examiner : Ron Schadron  
Art Unit : 1644  
For : **INDUCING ANTIBODY RESPONSE AGAINST SELF-PROTEINS  
WITH THE AID OF FOREIGN T-CELL EPITOPES**  
745 Fifth Avenue, New York, New York 10151

**DECLARATION OF ROLF M. ZINKERNAGEL, PhD**

Assistant Commissioner for Patents

Washington, D.C. 20231

Dear Sir:

**ROLF M. ZINKERNAGEL declares and says that:**

1. I am familiar with the subject matter of the above-captioned application (the present application) as: I am informed that a concurrently-filed Amendment presents claims as reproduced below or substantially as reproduced below, after my signature, which I have read and understood; I have been informed that the Examiner has indicated that Dr. Paul Travers - a previous declarant in the present application - is not a person knowledgeable in the field of immunology or more specifically in the field relating to immunosuppression; and I have been informed that the Examiner, in rejecting claims, has indicated that it would have been obvious for an immunologist to substitute suppressor epitopes in self-proteins with foreign T-helper epitopes or that an immunologist could have substituted suppressor epitopes in self proteins with T-helper epitopes. My *Curriculum vitae*, which is publicly available at *inter alia* <http://www.nobel.se/medicine/laureates/1996/zinkernagel-cv.html>, is attached and incorporated herein by reference. Among other accomplishments, I was awarded the Nobel Prize in Physiology or Medicine with Peter C. Doherty, PhD, in 1996, for our discoveries in concerning the specificity of the cell mediated immune defense. Accordingly, I respectfully submit that I am well qualified to speak as to the present application and Dr. Travers' qualifications.



**DR. TRAVERS IS WELL KNOWN AS BEING QUITE  
KNOWLEDGEABLE IN THE FIELD OF IMMUNOLOGY  
AND IN THE FIELD OF IMMUNOSUPPRESSION**

2. As mentioned above, I am informed that the Examiner has indicated that Dr. Paul Travers is not a person knowledgeable in the field of immunology, or more specifically, in the field relating to immunosuppression. I respectfully disagree with the Examiner's opinion of Dr. Paul Travers.

3. Dr. Paul Travers is the prominent author of the preeminent textbook Janeway/Travers Immunology. This textbook is probably the most widely known and recognized textbook in immunology in the world. Hence, Dr. Paul Travers is an internationally acknowledged and recognized immunologist. Dr. Travers, in my opinion, has a profound knowledge of the basic functioning of the immune system, especially as reflected by Dr. Travers being the prominent author of the preeminent textbook Janeway/Travers Immunology – again probably the most widely known and recognized textbook in immunology in the entire world.

4. Accordingly, I respectfully submit that no one in the field of immunology would or could reasonably concur with the Examiner's opinion that Dr. Paul Travers is not a person knowledgeable in the field of immunology, or more specifically, in the field relating to immunosuppression; and, I respectfully disagree with this opinion of Dr. Paul Travers by the Examiner.

**THE EXAMINER'S HYPOTHETICAL  
SUBSTITUTION OF SUPPRESSOR  
EPITOPES IS NOT AND WAS NOT POSSIBLE**

5. I am also informed that the Examiner has indicated that it would have been obvious for an immunologist to substitute suppressor epitopes in self-proteins with foreign T-helper epitopes. As a person clearly skilled in the field of immunology, and indeed recognized as an expert in the field of immunology, I respectfully submit, based on my education, training and experience, that the Examiner's hypothetical substitution of suppressor epitopes in self-proteins with foreign T-helper epitopes, is today not possible, and was not possible at the August 26, 1993 effective filing date of the present application; and therefore, that it could not have been obvious for an immunologist to substitute suppressor epitopes in self-proteins with foreign T-helper epitopes.



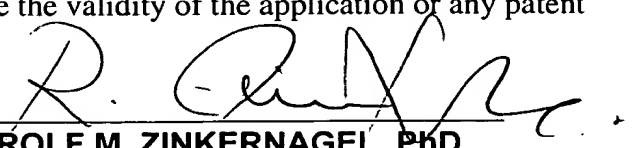
6. Simply, as a person clearly skilled in the field of immunology, and indeed recognized as an expert in the field of immunology, based on my education, training and experience, I fail to see how the Examiner's hypothetical substitution of suppressor epitopes could be possible today or could have been possible at the August 26, 1993 effective filing date of the present application. Even today, it is highly controversial whether there exists such a thing as suppressor epitopes; but, more importantly, there is, to the best of my knowledge, no known method of positively identifying such suppressor epitopes. And at the August 26, 1993 effective filing date of the present application, to the best of my knowledge, there was no known method of positively identifying such suppressor epitopes. Consequently it is not possible today – and was not possible at the August 26, 1993 effective filing date of the present application - to devise any strategy for substituting suppressor epitopes with foreign T-helper epitopes. Clearly, if one skilled in the art could not and cannot positively identify suppressor epitopes in self-proteins (whose existence is still a matter of debate in the art), based on my education, training and experience, there is no way and was no way for the skilled artisan to perform the hypothetical substitution postulated by the Examiner.

7. Thus, it was not obvious and is not obvious for an immunologist to substitute suppressor epitopes in self-proteins with foreign T-helper epitopes, contrary to the Examiner's hypothesis that it would have been obvious to substitute suppressor epitopes in self-proteins with foreign T-helper epitopes, with which, based on my education, training and experience, I respectfully disagree.

8. I further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and, that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful statements may jeopardize the validity of the application or any patent issued thereon.

Dated: June 8, 2002

By:

  
ROLF M. ZINKERNAGEL, PhD

**CLAIMS UNDERSTOOD TO BE ADDED OR SUBSTANTIALLY ADDED**

--56. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that

animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein;

whereby, the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

57. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein;

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

58. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving

tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least four amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

59. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least ten amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

60. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least fifteen amino acids on each side of the peptide fragment,

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

61. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken.

62. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by being detoxified and by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

63. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least four amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

64. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least ten amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

65. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving

secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least fifteen amino acids on each side of the peptide fragment,

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

66. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken.

67. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by being detoxified and by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

68. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising;

preparing different modified self-proteins, wherein:

each modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal,

said substitution preserving tertiary structure of the self-protein, and

the different modified self-proteins differ from each other with respect to the position of the at least one immunodominant T-cell epitope;

ascertaining which of the different modified self-proteins elicits a desired specific neutralizing effect and thereby ascertaining a desired modified self protein; and

administering to the animal, an immunologically effective amount of the desired modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein, and,

the desired modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

69. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, comprising;

preparing different modified self-proteins, wherein:

each modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal,

said substitution preserving secondary and tertiary structure of the self-protein, and

the different modified self-proteins differ from each other with respect to the position of the at least one immunodominant T-cell epitope;

ascertaining which of the different modified self-proteins elicits a desired specific neutralizing effect and thereby ascertaining a desired modified self protein; and

administering to the animal, an immunologically effective amount of the desired modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein, and,

the desired modified self-protein elicits antibodies that are against the self-protein, and B-cell autotolerance to the self-protein is broken.

70. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

a. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein;

whereby, the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

b. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

c. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least four amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

d. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least ten amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

e. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least fifteen amino acids on each side of the peptide fragment,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier

protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

f. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken; or,

g. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by being detoxified and by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

h. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution

preserving flanking regions comprising at least four amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

i. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self-protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least ten amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

j. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self-protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least fifteen amino acids on each side of the peptide fragment,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

k. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken; or,

l. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by being detoxified and by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

m. preparing different modified self-proteins, wherein:

each modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal,

said substitution preserving tertiary structure of the self-protein, and

the different modified self-proteins differ from each other with respect to the position of the at least one immunodominant T-cell epitope;

ascertaining which of the different modified self-proteins elicits a desired specific neutralizing effect and thereby ascertaining a desired modified self protein; and

administering to the animal, an immunologically effective amount of the desired modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein, and,

the desired modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

n. preparing different modified self-proteins, wherein:

each modified self-protein is modified, in comparison to the self-protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal,

said substitution preserving secondary and tertiary structure of the self-protein, and

the different modified self-proteins differ from each other with respect to the position of the at least one immunodominant T-cell epitope;

ascertaining which of the different modified self-proteins elicits a desired specific neutralizing effect and thereby ascertaining a desired modified self protein; and

administering to the animal, an immunologically effective amount of the desired modified self-protein, wherein:

the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein, and,

the desired modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken.

71. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, inducing antibody production in the animal against the self-protein of that animal, and eliciting an immune response in the animal which includes an MHC class II immune response as to an immunodominant T-cell epitope which is foreign to the animal and an

autoantibody response in other MHC-haplotypes, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

a. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing the immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken; or

b. the self-protein is normally autotolerated by the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing the immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken.

72. (New) A method for breaking B-cell autotolerance in an animal to a self-protein of that animal, and inducing antibody production in the animal against the self-protein of that animal, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, comprising administering to the animal, an immunologically effective amount of at least one modified self-protein, wherein:

a. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide

containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein;

whereby, the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

b. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

c. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least four amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

d. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide

containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least ten amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

e. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least fifteen amino acids on each side of the peptide fragment,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

f. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken; or,

g. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by being detoxified and by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self- protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

h. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least four amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self- protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

i. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least ten amino acids on each side of the peptide fragment;

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

j. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein, and said substitution preserving flanking regions comprising at least fifteen amino acids on each side of the peptide fragment,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

k. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes; and, the modified self-protein elicits an immune response in the animal which includes an MHC class II immune response as to the immunodominant T-cell epitope and an autoantibody response in other MHC-haplotypes, and B-cell autotolerance to the self-protein is broken; or,

l. the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein; and,

the modified self-protein is modified, in comparison to the self- protein, by being detoxified and by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal, said substitution preserving secondary and tertiary structure of the self-protein,

whereby the modified self-protein elicits antibodies that are against the self- protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

m. preparing different modified self-proteins, wherein:

each modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal,

said substitution preserving tertiary structure of the self-protein, and

the different modified self-proteins differ from each other with respect to the position of the at least one immunodominant T-cell epitope;

ascertaining which of the different modified self-proteins elicits a desired specific neutralizing effect and thereby ascertaining a desired modified self protein; and

administering to the animal, an immunologically effective amount of the desired modified self-protein, wherein:

the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein, and,

the desired modified self-protein elicits antibodies that are against the self- protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken; or,

n. preparing different modified self-proteins, wherein:

each modified self-protein is modified, in comparison to the self- protein, by containing a substitution of at least one peptide fragment of the self-protein with a peptide containing at least one immunodominant T-cell epitope which is foreign to the animal,

said substitution preserving secondary and tertiary structure of the self- protein, and

the different modified self-proteins differ from each other with respect to the position of the at least one immunodominant T-cell epitope;

ascertaining which of the different modified self-proteins elicits a desired specific neutralizing effect and thereby ascertaining a desired modified self protein; and

administering to the animal, an immunologically effective amount of the desired modified self-protein, wherein:

the self-protein is normally non-immunogenic in the animal and there is normally B-cell autotolerance by the animal to the self-protein, and,

the desired modified self-protein elicits antibodies that are against the self-protein, earlier and in higher titres, in comparison to the self-protein conjugated to a carrier protein or peptide containing T-cell epitopes, and B-cell autotolerance to the self-protein is broken.

73. (New) The method of any one of claims 56-72 wherein the modified self-protein is a recombinant modified self-protein.

74. (New) The method of any one of claims 56-72 wherein the self-protein is tumor necrosis factor alpha (TNF- $\alpha$ ), tumor necrosis factor beta (TNF- $\beta$ ), gamma interferon ( $\gamma$ -interferon), interleukin 1 (IL-1) or immune globulin (IgE).

75. (New) The method of claim 73 wherein the self-protein is tumor necrosis factor alpha (TNF- $\alpha$ ), tumor necrosis factor beta (TNF- $\beta$ ), gamma interferon ( $\gamma$ -interferon), interleukin 1 (IL-1) or immune globulin (IgE).

76. (New) The method of any one of claims 56-72 wherein the administering includes administering an adjuvant.

77. (New) The method of claim 76 wherein the adjuvant comprises calcium phosphate, saponin, quil A or a biodegradable polymer.

78. (New) The method of claim 73 wherein the administering includes an adjuvant.

79. (New) The method of claim 75 wherein the administering includes an adjuvant.--

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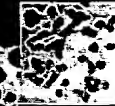
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Publications  
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**Education**

1962 Mathematisch-Naturwissenschaftliches Gymnasium,  
Basel, Matura

1962– 1968 Faculty of Medicine, University of Basel

1967– 1968 Course in Tropical Medicine, Tropical Institute, University  
of Basel

1968 National Board Examination

1968 E.C.F.M.G. (USA)

1970 M.D. Thesis

1971 Postgraduate course in Experimental Medicine, Faculty of  
Medicine, University of Zürich

1975 Ph.D.Thesis, Australian National University, Canberra,  
Australia

**Professional Record**

1966 Externship, Glen Cove Community Hospital, Glen Cove,  
Long Island, N.Y., USA

1969 Internship, Surgical Department, Clara-Spital, affiliated to the  
Facility of Medicine, University of Basel

1969 – 1970 Postdoctoral Fellow, Laboratory for Electron

Microscopy, Institute of Anatomy, University of Basel  
1971 – 1973 Postdoctoral Fellow, Institute of Biochemistry,  
University of Lausanne, Switzerland  
1973 – 1975 Visiting Fellow, Department of Microbiology,  
The John Curtin School of Medical Research, Australian  
National University, Canberra, Australia  
1975 – 1976 Associate (Assistant Professor), Department of  
Immunopathology, Research Institute of Scripps Clinic,  
La Jolla, California  
1976 – 1979 Associate Member (Associate Professor),  
Department of Immunopathology, Scripps Clinic and  
Research Foundation, La Jolla, California  
1977 – 1979 Adjunct Associate Professor, Department of  
Pathology, UCSD  
1979 Member (Full Professor), Department of Immunopathology,  
Scripps Clinic and Research Foundation  
1979 – 1988 Associate Professor, Department of Pathology,  
University of Zürich, University Hospital, Zürich  
1988 – 1992 Full Professor, Department of Pathology, University  
of Zürich, University Hospital, Zürich  
1992 – Head, Institute of Experimental Immunology, Zürich

**Honorary and Professional Organizations**

1971 – Swiss Society of Allergy and Immunology (President  
1993 – Honorary member 1996)  
1973 – 1975 Australian Society for Immunology (Honorary  
member 1996)  
1977 – American Association of Immunologists (Honorary  
member 1993)  
1977 – American Association of Pathologists  
1978 – Scandinavian Society of Immunology (Honorary member  
1978)  
1980 – Société Française d'Immunologie (Honorary member 1980)  
1980 – Swiss Society of Microbiology  
1981 – Swiss Society of Pathology  
1984 – EMBO European Molecular Biology Organization  
1987 – Swiss Association for the Study of the Liver  
1989 – Swiss Society of Cell and Molecular Biology  
1989 – Academia Europea  
1990 – Gesellschaft für Immunologie  
1990 – International Society for Antiviral Research  
1990 – ENI European Network of Immunological Institutions  
1990 – Deutsche Gesellschaft für Virologie  
1991 – Deutsche Gesellschaft für Immunologie  
1992 – The Delphinium Society  
1994 – Schweizerische Akademie der Medizinischen  
Wissenschaften  
1994 – Deutsche Akademie der Naturforscher Leopoldina  
1996 – American Academy of Microbiology, Fellow  
1996 – US National Academy of Sciences, Foreign Fellow  
1996 – Australian Academy of Sciences, Foreign Fellow  
1998 – American Academy of Arts and Sciences, Foreign Fellow  
1998 – Royal Society, Foreign Fellow  
1998 – Academie Royale de Medicine de Belgique, Foreign Fellow  
1998 – Berlin-Brandenburgische Akademie der Wissenschaften  
1998 – Foundation Gen Suisse  
1998 – WIF World Innovation Foundation, Honorary Member  
1998 – ECEAR European Conference on Experimental AIDS  
Research  
1999 – FMH Verband der Schweizer Ärzte und Ärztinnen  
2000 – Schweizer Wissenschafts- und Technologierat  
2001 – Patronage Committee Foundation Swiss Bridge

**Editorial Board**

1976 – 1988 Experimental Cell Biology  
1977 – Immunogenetics  
1978 – 1984 Parasite Immunology

1978 – 1980 Journal of Immunology  
1979 – 1989 Thymus  
1980 – Zeitschrift für Immunologie-Immunobiology  
1980 – 1988 Antiviral Research  
1981 – European Journal of Immunology (Executive Committee 1994)  
1981 – Journal of Environmental Pathology Toxicology & Oncology  
1981 – 1984 Journal of Experimental Medicine  
1981 – 1983 Current Topics in Microbiology and Immunology  
1982 – Scandinavian Journal of Immunology  
1983 – Cellular Immunology  
1983 – International Journal of Microbiology  
1987 – 1989 Europ.Molecular Biology Organization Journal  
1987 – European Journal of Clinical Investigation  
1988 – 1991 Journal of Autoimmunity  
1988 – 1991 Clinical Immunology and Immunopathology  
1988 – International Immunology  
1988 – 2000 Urologia Internationalis  
1989 – Annual Review of Immunology  
1991 – International Review of Experimental Pathology  
1991 – International Journal of Clinical & Laboratory Research  
1992 – Immunology today  
1992 – Immunology and Cell Biology  
1994 – Immunity  
1994 – Viral Immunology  
1994 – 2000 Virology  
1995 – Immunological Reviews  
1996 – Cell and Tissue Research  
1997 – Seminars in Immunopathology  
1997 – Current Opinion in Microbiology  
1998 – International Journal of Molecular Medicine  
1998 – History and Philosophy of the Life Sciences  
2000 – Cytokine

#### Honours

1981 Cloëtta Stiftung (Zürich)  
1982 Jung Stiftung (Hamburg)  
1983 Paul Ehrlich Preis (Frankfurt)  
1985 Mack-Forster Preis (Europ.Ass.Clin.Inv.)  
1986 Gairdner Foundation (Toronto)  
1987 Institute for Cancer Research (New York)  
1988 Louis Jeantet Foundation (Geneva)  
1988 Naegeli Stiftung (Zürich)  
1992 Christoforo Colombo Award (Genova)  
1995 Lasker Award (New York)  
1996 Honorary Dr. h.c., University of Liège  
1996 Honorary Dr. h.c., Australian National University, Canberra  
1996 Nobel Prize for Medicine or Physiology  
1997 Honorary Dr. h.c., University of Oslo  
1997 Honorary Dr. h.c., University of Quebec  
1997 Honorary Dr. h.c., University of Genova  
1997 Drew-Novartis Award  
1997 Reichstein Medaille (Zürich)  
1998 Honorary Dr. h.c., Latvian University, Riga  
1998 Honorary Dr. h.c., Agricultural University of Warsaw  
1999 Honorary Companion in the General Division of The Order of Australia  
1999 Member of the Order pour le mérite for Sciences and Arts (Germany)  
1999 Honorary Dr. h.c., University of Basel, Switzerland  
2000 Honorary Dr. h.c., University of Montréal, Canada  
2000 Honorary Dr. h.c., University of Buenos Aires, Argentina  
2000 Honorary Dr. h.c., Medical Academy of the University of Warsaw  
2000 Honorary Dr. h.c., Medical University of Odessa

#### Scientific Advisory Board

1977 – Temporary Adviser National Science Foundation

Washington

1979 – 1980 Study section NIH Virology  
1980 NIH-Task Force Immunology  
1981 – 1983 Gutachter Sonderforschungsbereich 107  
    «Vollzugsmechanismus der Immunreaktion», Mainz, DFG  
1981 – 1983 Ministère de la recherche et de l'Industrie Action  
    «Régulations en Immunologie et Immunopathologie» Paris  
1982 – 1983 Schwerpunktprogramm Histokompatibilitätsgen-  
    komplex Deutsche Forschungsgemeinschaft  
1982 – 1986 Basel Institut für Immunologie, Swiss Scientific  
    Advisory Board  
1982 – 1991 Zentrum für Lehre und Forschung, Universität Basel,  
    Advisory Board  
1985 – 1989 Scientific Advising Group of Experts in Vaccine  
    Development WHO  
1986 – 1991 Scientific Advisory Board, Max Planck-Institute for  
    Biology, Tübingen  
1986 – 1988 Biogen, Advisor  
1988 – Cancer Research Institute (Scientific Advisory Council)  
1988 – Sandoz Prize for Immunology Committee  
1989 – Schweiz. Institut für Allergie und Asthmaforschung, Davos  
1989 – Academia Europaea, London  
1990 – 1992 Founding Committee Max Planck - Institute of  
    Infectiology  
1991 – Scuola Superiore d'Immunologia, Napoli  
1991 – 1993 EMBO Council  
1992 – 1996 Biozentrum der Universität Basel  
1992 – 2000 Marcel Benoit Preis Kommittee  
1992 – Universität Basel, Biozentrum  
1996 – 2000 Jung-Stiftung für Wissenschaft und Forschung,  
    Hamburg (Committee member)  
1997 – Fondation pour le Recherche sur le Vieillissement, Genève  
1998 – 2000 Zurich University Association  
1998 – Foundation Science et Cité, Bern  
1998 – 2000 WIF World Innovation Foundation, Huddersfield GB  
1998 – Swiss Bridge Foundation, Zürich  
1999 – Academy of Cancer Immunology, New York  
1999 – Biomedical Research and Study Centre, Riga  
1999 – SFO - Foundation to support Organ Donation, Zürich

Special lectures

1979 The Kinyoun Lecture NIH  
1980 Wellcome visiting Professorship, Denver  
1980 Special University of London Lecture  
1982 Campbell Memorial Lecture, Asilomar  
1983 A. v. Graefe Lecture, Berlin  
1983 Armauer Hansen Memorial Lecture, Addis Abeba  
1993 Grabar Lecture, French Society of Immunology  
1994 Harvey Lecture, New York  
1986 Peter Gorer Lecture, British Society of Immunology  
1997 Felix-Hoppe-Seyler Lecture  
1997 Landsteiner Lecture, Frankfurt  
1998 Hubert-Bloch Lecture, Basel  
1998 Leopold G. Koss Lecture, Bern  
1998 Wolfgang-Pauli Lecture, Zürich  
2000 Meyenburg Lecture, Heidelberg

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### **Bibliography Rolf Zinkernagel**

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5. Frei, P.C., Erard, P., and Zinkernagel, R.M.  
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